#### (19) World Intellectual Property Organization International Bureau





(43) International Publication Date 17 August 2006 (17.08.2006)

## (10) International Publication Number WO 2006/084950 A1

(51) International Patent Classification: C22B 15/00 (2006.01) C22B 3/20 (2006.01)

(21) International Application Number:

PCT/FI2006/000046

(22) International Filing Date:

13 February 2006 (13.02.2006)

(25) Filing Language:

Finnish

(26) Publication Language:

English

(30) Priority Data: 20050160

14 February 2005 (14.02.2005)

(71) Applicant (for all designated States except US): OUT-OKUMPU TECHNOLOGY OY [FI/FI]; Riihitontuntie 7, FI-02200 Espoo (FI).

(72) Inventors; and

- (75) Inventors/Applicants (for US only): HULTHOLM, Stig-Erik [FI/FI]; Sitomantie 1, FI-28360 Pori (FI). HEIMALA, Seppo [FI/FI]; Marjastajantie 9, FI-28220 Pori (FI). RITASALO, Teemu [FI/FI]; Käenkatu 1 B 9, FI-29200 Harjavalta (FI). RUONALA, Mikko [FI/FI]; Pajukuja 6, FI-02460 Kantvik (FI).
- (74) Agent: OUTOKUMPU TECHNOLOGY OY, IPR MANAGEMENT; P.O. Box 86, FI-02201 Espoo (FI).

- (81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, LY, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.
- (84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

#### Declarations under Rule 4.17:

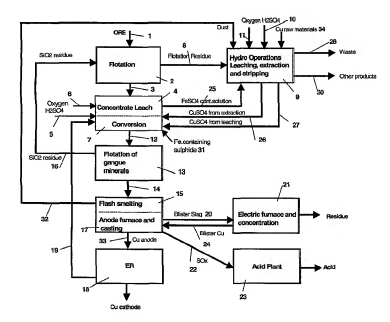
- as to applicant's entitlement to apply for and be granted a patent (Rule 4.17(ii))
- of inventorship (Rule 4.17(iv))

#### Published:

with international search report

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

(54) Title: METHOD FOR THE TREATMENT OF COPPER-BEARING MATERIALS



(57) Abstract: The present invention relates to a method for the treatment of copper-bearing materials, such as copper concentrates, so that the impurities and gangue minerals in the materials, such as silicates, are in effect removed completely.

## METHOD FOR THE TREATMENT OF COPPER-BEARING MATERIALS

The present invention relates to a method for the treatment of copper-bearing materials, such as copper concentrates, so that the impurities and gangue 5 minerals in the materials, such as silicates, are in effect removed completely.

Methods are known from WO patent applications 2005/007905, 2005/007902 and 2005/007901, in which copper, nickel and precious metals are effectively turned into sulphidic intermediate products with high copper content and sufficient caloric value using concentration and hydrometallurgical methods. This occurs in solution according to example reaction (1)

$$xCu^{2+} + yCuFeS_2 -> zCu_xS + yFeSO_4 + (y-z)H_2SO_4$$
 (1),

where the copper in solution is turned into copper sulphide form by conversion using iron-containing sulphide material. A conversion reaction occurring with the same kind of iron-containing sulphide material is used for example to recover nickel and precious materials from solution as well in nickel-copper matte leaching.

20

By exploiting the known methods described in the above-mentioned WO patent applications 2005/007905, 2005/007902 and 2005/007901, the iron content for instance of copper concentrate can be effectively reduced. However, some problems arise in these methods from time to time, because the copper concentrate obtained in conversion still contains gangue minerals, such as silicates. If this kind of iron-depleted copper concentrate is used for example in copper smelting, these silicates require the addition of iron, so that the slag can be made sufficiently liquid to get the slag out of the smelter at the desired temperature. As a result, the amount of copper smelting slag increases.

30

The purpose of the present invention is to eliminate the disadvantages of the prior art and to achieve an improved method for the treatment of copper-

bearing materials, such as copper concentrates, so that the impurities and gangue minerals in the materials, such as silicates, can in effect be removed completely. The essential features of the invention will be made apparent in the attached claims.

5

In the method according to the invention, conditions are applied to the copper-bearing material to be treated, such as copper concentrate, in the presence of firstly an acid, such as sulphuric acid, hydrochloric acid or an equivalent acid, and an oxygen-containing gas such as oxygen, air or oxygen-enriched air, in which the copper obtained in solution is turned by conversion with a material containing iron sulphide into copper sulphide form, using the principle reaction (1) presented above. The sulphuric acid - ferrous sulphate solution thus generated also contains the impurities of the leached copper-bearing material, such as arsenic, antimony, bismuth, uranium, zinc, nickel and cobalt contents.

15 With the copper sulphide formed by conversion the gangue minerals e.g. silicates, can be made to separate according to the invention by flotation and/or a method based on differences in specific weight.

In the flotation of the material containing copper sulphide obtained by conversion according to the invention, the flotation process is preferably carried out so that the copper sulphides and the precious metals with them in the copper sulphide containing material are frothed, so that the gangue minerals, such as silicates, are made to enter the residue of the flotation process by means of depressant chemicals. The flotation process may also be carried out so that the gangue minerals, such as silicates, are frothed and the copper-bearing sulphide material together with the precious metals is recovered in the residue of the flotation process by means of depressant chemicals. When proceeding in either way, specified electro-chemical potential or electro-chemical alternating current spectroscopic methods using mineral electrodes can be exploited advantageously in the flotation process. The separation of gangue minerals is improved and the iron content of the copper sulphide product of conversion is reduced beneficially when the conversion temperature

is maintained in a range between 170 - 260°C, preferably between 200 - 220°C.

If there is a substantial amount of chlorides and/or fluorides in the water used in the leaching or other treatment of copper-bearing material, not only gangue minerals such as silicates, but also halides, such as chlorides and/or fluorides for example, can be separated effectively from the usually coarse copper sulphide material obtained in conversion, the majority preferably even during the water separation of the copper sulphide material obtained in conversion.

This is possible due to the coarseness and thus excellent filterability and low residual moisture of the copper sulphide conversion product. The separation of the copper sulphide material and the halides can be intensified by performing filtration and/or washing of the copper sulphide material in reductive conditions, which are achieved using chemicals and/or an electric current.

15

The method according to the invention is very practicable for various copperbearing materials and many localised water supply and water quality problems.

The method according to the invention can be used when treating copper scrap, smelter dust and copper-zinc-lead ores in addition to copper-bearing ores. The last mentioned ores often include minerals containing precious metals such as arsenopyrite or pyrite, which can be converted to copper sulphide minerals, which contain said precious metals according to the the method. Furthermore, the method according to the invention allows the use of salt-containing water, e.g. seawater, as process water. These presuppose that process parameters obtained using known methods as such are exploited when regulating the process e.g. by using mineral electrodes and catalysts in process optimisation and the use of various leaching stages, such as heap leaching, for example to regulate the copper and acid balances in the overall process. In this way the object of the invention i.e. the separation of copper sulphide and gangue minerals can be intensified.

There is still some degree of copper and other valuable components in the silicate-containing fraction rich in SiO<sub>2</sub> obtained from the separation process carried out after conversion preferably by flotation. The copper and other valuable components in the SiO<sub>2</sub>-rich fraction can be recovered by leaching the fraction separately or during a leaching stage related to some other overall copper-bearing material process. The SiO<sub>2</sub>-rich fraction can also be returned to the flotation of the original copper-bearing ore. In addition a partial separation of the coarse sulphides can be performed if necessary on the SiO<sub>2</sub>-rich fraction from the post-conversion separation stage, before further treatment. Further, the gangue mineral product obtained from the post-conversion separation stage, the removal of gangue minerals, can be routed by means of mineral electrodes to electro-chemically controlled further processing, such as for instance leaching or flotation.

15 When the method of the invention is applied e.g. to slag from a copper smelting process, the copper content of the slag is first turned into sulphide form by conversion and the gangue mineral content e.g. the silicon dioxide (SiO<sub>2</sub>) content of the slag decreases in the flotation of the sulphide product obtained by conversion. When feeding the copper sulphide-containing flotation product with low SiO<sub>2</sub> content thus obtained back to a smelter that uses sulphidic copper raw material, the amount of slag exiting the smelter is in effect reduced by this amount.

Due to the low SiO<sub>2</sub> content, the slag obtained from the smelting of sulphidic copper-bearing material can be treated in several ways. The slag can advantageously be granulated for example and leached separately or together with the dust from the smelting of copper-bearing material and the SiO<sub>2</sub>-rich fraction obtained in post-conversion flotation with iron-containing sulphide according to the invention. In this way the copper and other valuable components remaining in connection with the SiO<sub>2</sub> fraction are made to dissolve at least partially, and thus are available for further processing without the copper requiring further slag treatment. The slag obtained from the smelting

of copper-bearing material can also be treated pyrometallurgically, preferably for example by smelting in an electric furnace. In this way any potential deficiency in soluble copper entering copper conversion is filled by leaching additional other copper-bearing raw materials to keep the copper conversion copper balance beneficial.

When exploiting the copper-bearing material treated with the method according to the invention in copper smelting, the Cu/Fe/SiO<sub>2</sub> ratios of the copper entering smelting and of the copper-rich conversion product obtained by conversion are 10 vitally important with regard to the final smelting result, and these Cu/Fe/SiO2 ratios can preferably be adjusted in conversion and in the related gangue mineral separation irrespective of essentially different copper-bearing raw materials, with mineral electrode measurements, for instance using electrochemical potential. The sulphur balance prevailing in the total process 15 must also be taken into account in mineral electrode measurements, in which the sub-components include notably S2- ions, elemental sulphur S0 and sulphate sulphur SO<sub>4</sub><sup>2</sup>. When using the method of the invention, whereby the essentially copper sulphide free phase of the gangue minerals can be fed onwards to e.g. copper smelting, and the SiO2-containing fraction of the 20 gangue minerals such as silicates can be treated to remove the copper remaining in the SiO<sub>2</sub>-containing fraction effectively, the copper yield of the overall process can be made extremely high in the final copper-containing product such as raw copper, optimally above 99 %.

When the method of the invention is applied to the smelting of copper-bearing material, the leaching of the material being processed, the conversion of copper into sulphide, and the separation of gangue minerals such as silicates from the sulphidic conversion product, can also be carried out completely separately from the smelting process, even in geographical terms. That which is transported in this case between gangue mineral separation and copper recovery is usually copper sulphide concentrate containing 65 – 77% copper.

The invention is described below in more detail with reference to the appended drawings, in which

- Fig 1 presents a flow chart of one preferred embodiment of the invention applied to copper smelting,
- 5 Fig 2 presents the results of the copper and SiO<sub>2</sub> fraction yield correlation in post-conversion flotation, and
  - Fig 3 shows the results of the copper and SiO<sub>2</sub> fraction content ratio in post-conversion flotation.
- 10 According to Figure 1, ore 1 containing copper sulphide and possibly other valuable components is routed to flotation 2 to obtain copper sulphide concentrate 3 that contains the valuable components. The copper sulphide concentrate obtained from flotation is routed onwards to leaching 4. Leaching 4 is performed in the presence of sulphuric acid 5 and an oxidising reagent, such 15 as oxygen 6, preferably at a temperature of 220°C, and the copper in the copper sulphide concentrate dissolves in leaching 4 into Cu<sup>+</sup>/Cu<sup>++</sup> ions. After leaching 4, the dissolved copper undergoes conversion 7 at a temperature of 220°C in the presence of material containing iron sulphide 31 according to principal reaction (1) in order to obtain copper sulphide product, ferrous sulphate and sulphuric acid. This solution of sulphuric acid and ferrous sulphate also contains the impurities in the leached copper sulphate concentrate, such as arsenic, uranium, zinc, nickel and cobalt contents.

The flow sheet shown in Figure 1 also describes the passage of the flotation residue 8 obtained from the copper-bearing ore flotation stage 2 to hydrometallurgical treatment 9, where leaching in the presence of an oxygen-containing reagent 11 is performed on the flotation residue, followed by extraction and stripping of the solution obtained with sulphuric acid 10. Since the ferrous salt-oxygen solution 25 from conversion 7 containing impurities can advantageously be routed to this hydrometallurgical treatment 9, this treatment 9 can function as a base for the recovery or removal of impurities in copper-bearing raw material, such as arsenic entering the waste 28, and other products

30, such as uranium, zinc, lead, nickel and cobalt. Furthermore, the copper sulphate solutions 26 and 27 that contain sulphuric acid obtained from the leaching and extraction stage of hydrometallurgical treatment 9 can also be fed to the leaching stage 4 in addition to the conversion stage 7.

5

The copper sulphide product 12 obtained in conversion 7 using copper sulphate and sulphuric acid is routed according to the invention to the removal of gangue materials, such as silicates, by flotation 13. The copper sulphide product 14 obtained from flotation 13 is taken to the flash smelter 15 for raw copper 10 fabrication. The flotation residue 16 obtained from flotation 13, which contains gangue materials, such as silicates, is returned to the original flotation 2 of copper-bearing ore 1.

The raw copper 33 obtained from the flash smelter 15 is routed in molten form to the anode furnace 17 and also on to be cast from molten raw copper as anodes suitable for the electrolytic refining 18 of copper. The impure copper sulphate solution 19 containing sulphuric acid, obtained from electrolytic refining 18, is recirculated back to copper sulphide concentrate leaching 4, whereas cathodes are formed from the copper, and are used as raw material for the further processing of copper. Flash smelter dust 32 can be returned for example to hydrometallurgical treatment 9, into which other copper-containing raw materials 34 can also be fed if necessary.

The slag 20 obtained from the flash smelter 15 is routed to an electric furnace 25 21 and on to slag concentration, from where the copper-rich phase 24 obtained is conveyed back to the smelter 15 to be re-smelted. The exhaust gases 22 from the smelter 15 are routed to the sulphuric acid plant 23, and the sulphuric acid from there can be used in copper concentrate leaching 4.

## **EXAMPLE**

To test the method of the invention, the four separate tests mentioned in Tables 1 and 2 were carried out first, where conversion was performed on materials containing copper-containing copper minerals, such as chalcopyrite (CuFeS<sub>2</sub>), 5 bornite (Cu<sub>5</sub>FeS<sub>4</sub>) and digenite (Cu<sub>2-x</sub>S), and gangue minerals such as silicates and iron oxides, in the presence of iron-containing sulphide, such as e.g. pyrite (FeS<sub>2</sub>) or arsenopyrite (FeAsS). The total SiO<sub>2</sub> content of the gangue minerals in the materials processed was 5.2 % by weight.

- 10 In tests 1 − 3, the copper solution was brought to conversion directly from copper-bearing material leaching and partially concentrated by means of liquid-liquid extraction of copper after heap leaching. In test 4, a combined leaching and conversion test was carried out, in which the first interval from 0 − 30 min was the leaching stage and the 30 − 60 min interval was the conversion stage.
- 15 The process stages were monitored and regulated using mineral and platinum electrodes and, for instance, the energy levels of the electrodes in the conversion stage were kept between +300 +370 mV vs. AgCl/Ag.

Test	°c	Fe, % time, min				
		0	30	60		
1	160	14.5	8.3	7.2		
2	190	14.5	7.4	5.0		
2 3 4	220	14.5	1.85	1.8		
4	220	14.5	6.1	2.6		

## 20 Table 1: Fe content as a function of time

Test	°c	Cu, g/l time, min			
		0	30	60	
1	160	59	38.7	27.6	
2	190	59	10.8	4.1	
3	220 220	41	0	0	
4	220	9.5	0.45	0.008	

Table 2: Cu content as a function of time

According to Tables 1 and 2 the conversion temperature in tests 1 – 3 was 160°C, 190°C and 220°C respectively, whereas in test 4 both the leaching 5 temperature and the conversion temperature was 220°C. Table 1 shows the change of iron content in the solids during conversion as a function of time and Table 2 shows the copper content of the solution as a function of time. In tests 1 – 3 the SiO<sub>2</sub> content of the solids remained almost constant, whereas in test 4 the final SiO<sub>2</sub> content almost doubled.

10

The method according to the invention was applied to the final residue from test 3, which was floated to recover the copper sulphide and precious metal contents still in the final sediment. The results of flotation are given in Figures 2 and 3. Figure 2 shows the yield dependency of the copper and SiO<sub>2</sub> fractions 15 and Figure 3 shows the content ratio of the copper and SiO<sub>2</sub> fractions. The terms used in Figs. 2 and 3 mean the following: RC rough concentrate, SC scavenger concentrate, CC1 concentrate from the first cycle and CC2 concentrate from the second cycle. Flotation was performed so that the material was first pre-frothed, then scavenging was performed on the rough concentrate. In this way repeat flotation was performed on the resulting concentrate in two stages.

In Figure 2 the yield dependency of the copper and SiO<sub>2</sub> fractions is described in a graph, where the x-axis shows the copper yield (wt.%) and the y-axis the yield of SiO<sub>2</sub> (wt.%). In Figure 2 the starting point (feed) is that the yield from each fraction is 100 wt.%. The graph shows that after rough concentration and scavenger concentration the SiO<sub>2</sub> yield has fallen to about 40% of the starting point, whereas the copper yield is about 95 % of the starting point. After two repeat concentration cycles (CC2) there is only about 10% of the original yield of SiO<sub>2</sub>, which means that about 90% of the SiO<sub>2</sub> content has been removed. On the other hand, there is still about 63% of the original copper content.

In Figure 3 the content dependency of the copper and SiO<sub>2</sub> fractions is depicted in a graph, where the x-axis shows the copper content (wt.%) and the y-axis the SiO<sub>2</sub> content (wt.%). At the starting point of flotation (feed) the SiO<sub>2</sub> content is 5.2 wt.% and the copper content about 61.7 wt.%. After the two repeat flotation stages the SiO<sub>2</sub> content has fallen to a value of 1.0 wt.%, whereas the copper content has risen to a value of 67.5 wt.%

#### PATENT CLAIMS

- A method for treating copper-bearing material, in which the copper-bearing material is leached (4) and conversion (7) is performed on the leaching product
   in the presence of material containing iron sulphide (31) in order to obtain copper sulphide (12), characterised in that gangue mineral separation (13) is carried out on the copper sulphide (12) attained in conversion before the further processing of said copper sulphide (12).
- 10 2. A method according to claim 1, **characterised in that** the separation of gangue minerals (13) is carried out by flotation.
- A method according to claim 2, characterised in that the valuable components of the copper sulphide are floated apart from the gangue minerals
   during gangue mineral separation (13).
  - 4. A method according to claim 2, **characterised in that** the gangue minerals are floated apart from the valuable components of the copper sulphide during gangue mineral separation (13).

20

- 5. A method according to claim 1, **characterised in that** separation of gangue minerals (13) is performed on the basis of the difference in specific weight.
- 6. A method according to any of the preceding claims, characterised in that, in order to improve the separation of gangue minerals (13), the copper sulphide conversion (7) preceding gangue mineral removal takes place at a temperature range of 170 260°C.
- 7. A method according to claim 6, **characterised in that**, in order to improve 30 the separation of gangue minerals (13), the copper sulphide conversion (7) preceding gangue mineral removal takes place preferably at a temperature range of 200 220°C.

8. A method according to any of the preceding claims, characterised in that the gangue mineral product (16) obtained from gangue mineral separation (13) is routed to the leaching (4) that precedes conversion (7).

5

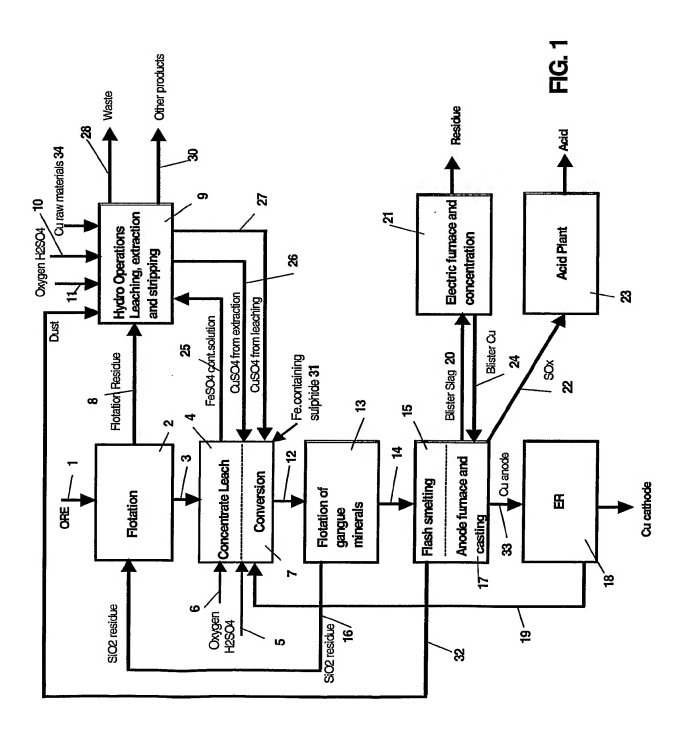
9. A method according to any of the preceding claims 1 - 7, **characterised in that** the gangue mineral product (16) obtained from gangue mineral separation (13) is routed to the concentration (2) of copper-bearing ore that precedes conversion.

10

10. A method according to any of the preceding claims 1 - 7, **characterised in that** the gangue mineral product (16) obtained from gangue mineral separation (13) is routed to further processing controlled electrochemically by means of mineral electrodes, such as e.g. leaching or flotation.

15

- 11. A method according to any of the preceding claims, **characterised in that** the copper sulphide product (14) obtained from gangue mineral separation (13) is routed to a smelter (15) for the fabrication of raw copper.
- 20 12. A method according to any of the preceding claims, **characterised in that** electrochemical potential measurement with mineral electrodes is exploited in the separation of gangue minerals (13).
- 13. A method according to any of the preceding claims, characterised in that25 an alternating current spectroscopic method is exploited in the separation of gangue minerals (13).
- 14. A method according to any of the preceding claims, characterised in that the separation of halides, such as chloride and/or fluoride, is performed in connection with the separation of water from the copper sulphide material (12) obtained in conversion (7).



SiO2 yield (%)

100
90
80
70
60
50
40
30
20
10
60
70
80
90
100
Cu yield (%)

FIG. 2

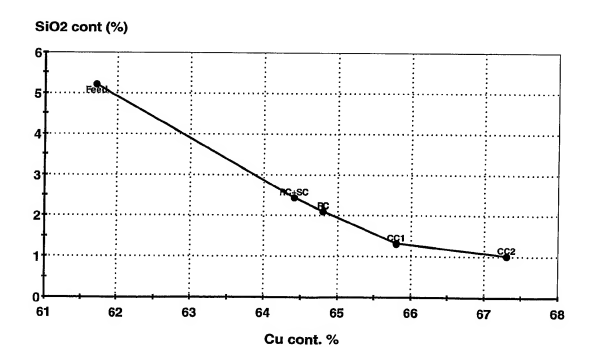


FIG. 3

#### INTERNATIONAL SEARCH REPORT

International application No. PCT/FI2006/000046

## A. CLASSIFICATION OF SUBJECT MATTER IPC: see extra sheet According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) IPC: C22B Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched SE,DK,FI,NO classes as above Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) EPO-INTERNAL, WPI DATA, PAJ C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Category\* Citation of document, with indication, where appropriate, of the relevant passages A WO 2005007902 A1 (OUTOKUMPU OYJ), 27 January 2005 1-14 (27.01.2005), figure 1, abstract WO 2005007901 A1 (OUTOKUMPU OYJ), 27 January 2005 1-14 A (27.01.2005), figure 1, abstract WO 2005007905 A1 (OUTOKUMPU OYJ), 27 January 2005 Α 1 - 14(27.01.2005), figure 1, abstract Α US 5616168 A (PHILIP J. GABB ET AL), 1 April 1997 1-14 (01.04.1997), figure 1, abstract X See patent family annex. Further documents are listed in the continuation of Box C. Special categories of cited documents: later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "A" document defining the general state of the art which is not considered to be of particular relevance earlier application or patent but published on or after the international filing date "E" document of particular relevance: the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) document of particular relevance: the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art document referring to an oral disclosure, use, exhibition or other document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report **3 0 -**05- 2006 22 May 2006 Name and mailing address of the ISA/ Authorized officer Swedish Patent Office Box 5055, S-102 42 STOCKHOLM Anna-Maj Magnusson/MP Facsimile No. +46 8 666 02 86 Telephone No. +46 8 782 25 00

Form PCT/ISA/210 (second sheet) (April 2005)

#### INTERNATIONAL SEARCH REPORT

International application No. PCT/FI2006/000046

# International patent classification (IPC)

C22B 15/00 (2006.01) C22B 3/20 (2006.01)

# Download your patent documents at www.prv.se

The cited patent documents can be downloaded at <a href="www.prv.se">www.prv.se</a> by following the links:

- In English/Searches and advisory services/Cited documents (service in English) or
- e-tjänster/anförda dokument(service in Swedish).

Use the application number as username. The password is **SJLDUSVIFW**.

Paper copies can be ordered at a cost of 50 SEK per copy from PRV InterPat (telephone number 08-782 28 85).

Cited literature, if any, will be enclosed in paper form.

Form PCT/ISA/210 (extra sheet) (April 2005)

# INTERNATIONAL SEARCH REPORT

Information on patent family members

04/03/2006

International application No. PCT/FI2006/000046

WO	2005007902	A1	27/01/2005	AU CA FI FI	2004257843 / 2530354 / 116070   20031083 /	Å B	27/01/2005 27/01/2005 15/09/2005 18/01/2005
WO	2005007901	A1	27/01/2005	AU CA FI FI	2004257844 / 2530355 / 116683   20031084 /	A B	27/01/2005 27/01/2005 31/01/2006 18/01/2005
WD	2005007905	A1	27/01/2005	AU FI FI	2004257842 / 116686   20031082 /	В	27/01/2005 31/01/2006 18/01/2005
US	5616168	A	01/04/1997	US AU CA EP ES WO	5443622 5407096 2216119 0815269 2167556 9631629	A A A,B T	22/08/1995 23/10/1996 10/10/1996 07/01/1998 16/05/2002 10/10/1996

Form PCT/ISA/210 (patent family annex) (April 2005)